FORM 6-K

SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

Report of Foreign Private Issuer

Pursuant to Rule 13a-16 or 15d-16 under the Securities Exchange Act of 1934

For the month of May 2009		
Commission File Number	0-16174	

TEVA PHARMACEUTICAL INDUSTRIES LIMITED

(Translation of registrant's name into English)

5 Basel Street, P.O. Box 3190 Petach Tikva 49131 Israel

(Address of principal executive offices)

Indicate by check mark whether the Form 40-F:	registrant files or will file annual r	eports under	cover of Form 20-F or
Form 20-F	X	Form 40-F	
Indicate by check mark if the registra Rule 101(b)(1):	ant is submitting the Form 6-K in	paper as perr	mitted by Regulation S-T
Indicate by check mark if the registra Rule 101(b)(7):	ant is submitting the Form 6-K in	paper as perr	mitted by Regulation S-T
Indicate by check mark whether by f hereby furnishing the information to Exchange Act of 1934.			
Yes		No	X
If "Yes" is marked, indicate below the 2(b): 82	ne file number assigned to the regi	strant in con	nection with Rule 12g(3)-



TEVA PHARMACEUTICAL INDUSTRIES LTD.

Contact: Elana Holzman Teva Pharmaceutical Industries Ltd.

Kevin Mannix Teva North America

972 (3) 926-7554 (215) 591-8912

For Immediate Release

Lonza and Teva Announce Receipt of Regulatory Approval

-- Joint Venture Formally Established --

Jerusalem, Israel, and Basel, Switzerland, May 14, 2009 – Teva Pharmaceutical Industries Ltd. (NASDAQ: TEVA) and Lonza Group Ltd. (SIX: LONN) today announced that the European Commission granted antitrust approval to the joint venture between the two companies, which was originally announced on January 20, 2009.

In their joint venture Teva and Lonza will cooperate to develop, manufacture and market a number of affordable, efficacious and safe generic equivalents of a selected portfolio of biologic pharmaceuticals.

Teva and Lonza have now completed all formalities related to the establishment of the joint venture.

About Teva

Teva Pharmaceutical Industries Ltd., headquartered in Israel, is among the top 20 pharmaceutical companies in the world and is the world's leading generic pharmaceutical company. The Company develops, manufactures and markets generic and innovative human pharmaceuticals and active pharmaceutical ingredients, as well as animal health pharmaceutical products. Over 80 percent of Teva's sales are in North America and Europe.

About Lonza

Lonza is one of the world's leading suppliers to the pharmaceutical, healthcare and life science industries. Its products and services span its customers' needs from research to final product manufacture. Lonza is the global leader in the production and support of active pharmaceutical ingredients both chemically as well as biotechnologically. Biopharmaceuticals are one of the key growth drivers of the pharmaceutical and biotechnology industries. Lonza has strong capabilities in large and small molecules, peptides, amino acids and niche bioproducts which play an important role in the development of novel medicines and healthcare products. Lonza is a leader in cell-based research, endotoxin detection and cell therapy manufacturing. Lonza is also a leading provider of value chemical and biotech ingredients to the nutrition, hygiene, preservation, agro and personal care markets.

Lonza is headquartered in Basel, Switzerland and is listed on the SWX Swiss Exchange. In 2008, Lonza had sales of CHF 2.937 billion. Further information can be found at www.lonza.com.

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For further information:

Teva

Website: www.tevapharm.com

Contacts: Elana Holzman

Teva Pharmaceutical Industries Ltd.

Tel. 972 (3) 926-7554

elana.holzman@teva.co.il

Kevin Mannix Teva North America Tel. (215) 591-8912

kevin.mannix@tevausa.com

Lonza

www.lonza.com

Media Relations

Dominik Werner Tel +41 61 316 8798

dominik.werner@lonza.com

Investor Relations

Alexandre Pasini Tel +41 61 316 8835

alexandre.pasini@lonza.com

Safe Harbor Statement under the U. S. Private Securities Litigation Reform Act of 1995:

Teva's Safe Harbor Statement under the U. S. Private Securities Litigation Reform Act of 1995: This release contains forward-looking statements, which express the current beliefs and expectations of management. Such statements are based on management's current beliefs and expectations and involve a number of known and unknown risks and uncertainties that could cause our future results, performance or achievements to differ significantly from the results, performance or achievements expressed or implied by such forward-looking statements. Important factors that could cause or contribute to such differences include risks relating to: our ability to successfully develop and commercialize additional pharmaceutical products, the introduction of competing generic equivalents, the extent to which we may obtain U.S. market exclusivity for certain of our new generic products and regulatory changes that may prevent us from utilizing exclusivity periods, potential liability for sales of generic products prior to a final resolution of outstanding patent litigation, including that relating to the generic versions of Neurontin®, Lotrel® and Protonix®, the current economic conditions, competition from brand-name companies that are under increased pressure to counter generic products, or competitors that seek to delay the introduction of generic products, the effects of competition on our innovative products, especially Copaxone® sales, dependence on the effectiveness of our patents and other protections for innovative products, especially Copaxone®, the impact of consolidation of our distributors and customers, the impact of pharmaceutical industry regulation and pending legislation that could affect the pharmaceutical industry, our ability to achieve expected results though our innovative R&D efforts, the difficulty of predicting U.S. Food and Drug Administration, European Medicines Agency and other regulatory authority approvals, the uncertainty surrounding the legislative and regulatory pathway for the registration and approval of biotechnology-based products, the regulatory environment and changes in the health policies and structures of various countries, supply interruptions or delays that could result from the complex manufacturing of our products and our global supply chain, our ability to successfully identify, consummate and integrate acquisitions, including the integration of Barr Pharmaceuticals, Inc., the potential exposure to product liability claims to the extent not covered by insurance, our exposure to fluctuations in currency, exchange and interest rates, significant operations worldwide that may be adversely affected by terrorism, political or economical instability or major hostilities, our ability to enter into patent litigation settlements and the intensified scrutiny by the U.S. government, the termination or expiration of governmental programs and tax benefits, impairment of intangible assets and goodwill, environmental risks, and other factors that are discussed in our Annual Report on Form 20-F and in our other filings with the U.S. Securities and Exchange Commission ("SEC").



Web Site: www.tevapharm.com Teva Pharmaceutical Industries Ltd.

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

TEVA PHARMACEUTICAL INDUSTRIES LIMITED (Registrant)

/s/ Eyal Desheh Name: Eyal Desheh By:

Title: Chief Financial Officer

Date: May 14 2009